# Short Read Workshop Day 4 Trimming, Mapping, IGV

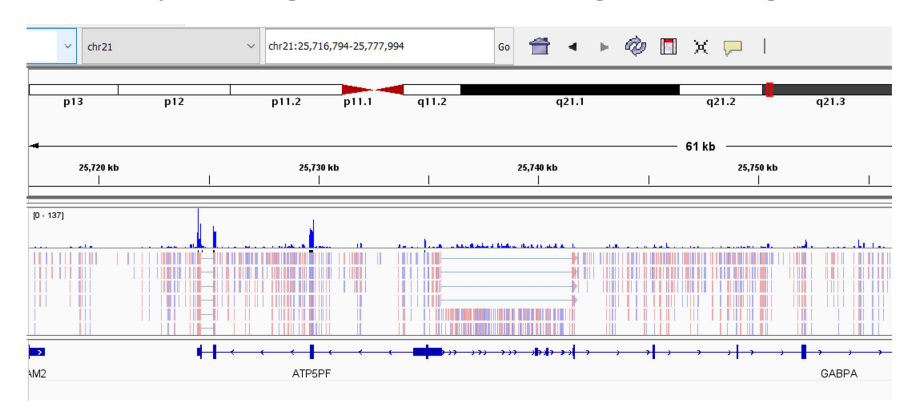
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### Day 4 overview

- Trimming fastq files
- Mapping fastq files
- More about mapped file formats
- Visualizing mapped files

# Goal of the Day

View sequencing data as reads aligned to a genome



50 base read:

TAGGCTAACTCTGTAGCCCCAGGTACCATGCATAATTGACCAGGATATAG

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Trimmomatic

40 base trimmed read: TAGGCTAACTCTGTAGCCCCAGGTACCATGCATAATTGAC

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HISAT2

Genome:

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HISAT2

Genome:

IGV

## Trimming fastq files with Trimmomatic

- Follow Trimmomatic worksheet to:
- Create Day4 directories
- Edit script to run Trimmomatic
  Input: fastq files with full-length reads
  Output: fastq files with trimmed/filtered reads

 Extra: Edit the d4\_trim\_qc.sbatch script to run pre-trim qc and compare the plots/stats to post-trim

# How do you trim polyA regions from both sides of reads?

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- Make a new fasta file with a polyA segment, or append to the Illumina adapter file, if writeable
- ILLUMINACLIP:<new fasta file>:2:30:10

### Mapping fastq files with HISat2

- Follow Mapping/IGV worksheet to:
- Rsync the mapping script
- Edit script and run HISAT2

Input: trimmed fastq files

Outputs: .sam, .bam, .sorted.bam, .bam.bai files

Visualize BAM file on the IGV web app

### Homework

Day 4 Homework – FASTQC, trimming, mapping, IGV

The assessment tomorrow will run many of the same steps as this homework. These steps are essential in ALL short read data processing.

# Variables – evaluating (calling)

#### \$variablename

echo \$a

grep \$gene\_name <filename>

trim\_script="\$filepath"/d4\_trim\_qc.sbatch OR

trim\_script=\${filepath}/d4\_trim\_qc.sbatch

wc \$filelist

#### Several ways of evaluating:

\$a

\${a}

"\$a"

These differ slightly, and you will see us use them all in scripts